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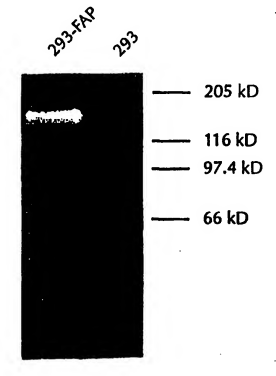
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(54) Title: ISOLATED DIMERIC FIBROBLAST ACTIVATION PROTEIN ALPHA, AND USES THEREOF

(57) Abstract

The invention involves dimeric forms of the protein known as fibroblast activation protein alfpha, or "FAPa" and its uses.



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ISOLATED DIMERIC FIBROBLAST ACTIVATION PROTEIN ALPHA, AND USES THEREOF

RELATED APPLICATION

This application is a continuation-in-part of Serial No. 08/230,491, filed April 20, 1994, now pending and incorporated by reference.

FIELD OF THE INVENTION

This invention relates to certain molecules associated with cancer tissues and reactive tumor stromal cells. particularly, it relates to fibroblast activation protein alpha ("FAP α " hereafter) molecules. A monomeric form of the molecule has previously been identified immunochemically, but nucleic acid molecules coding for it had not been isolated or cloned nor have dimers been identified. These, inter alia, are features of the invention. The monomeric protein has a molecular weight of from about 88 to about 95 kilodaltons as determined by SDS-PAGE of boiled samples. The dimer has a molecular weight of about 170 kilodaltons as determined by SDS-PAGE of unboiled samples. $FAP\alpha$ is characterized by a number of features and properties which are shared by and characteristic of membrane bound enzymes, suggesting very strongly that it, too, is a membrane bound enzyme. nucleic acid molecules, which are a key part of the invention, are useful both as probes for cells expressing FAPa, and as starting materials for recombinant production of the protein. The FAPa protein can then be used to produce monoclonal antibodies specific for the protein and are thus useful diagnostic agents themselves. They also have additional uses, including uses related to enzymatic functions, as described herein.

BACKGROUND AND PRIOR ART

The invasive growth of epithelial cancers is associated with characteristic cellular and molecular changes in the supporting stroma. For example, epithelial cancers induce the formation of tumor blood vessels, the recruitment of reactive tumor stromal fibroblasts, lymphoid and phagocytic

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infiltrates, the release of peptide mediators and proteolytic enzymes, and the production of an altered extracellular matrix See, e.g., Folkman, Adv. Cancer Res. 43: 175-203 (1985); Basset et al., Nature 348: 699-704 (1990); Denekamp et al., Cancer Metastasis Rev. 9: 267-282 (1990); Cullen et al., Cancer Res. 51: 4978-4985 (1991); Dvorak et al., Cancer Cells 3: 77-85 (1991); Liotta et al., Cancer Res. 51: 5054s-5059s (1991); Garin-Chesa et al., J. Histochem. Cytochem. 37: 1767-1776 (1989). A highly consistent molecular trait of the stroma in several common histologic types of epithelial cancers is induction of the fibroblast activation protein (FAP α), a cell surface glycoprotein with an observed M_r of 95,000 originally discovered with a monoclonal antibody, mAb F19, raised against proliferating cultured fibroblasts. Rettig et al., Cancer Res. 46: 6406-6412 (1986); Rettig et al., Proc. Natl. Acad. Sci. USA 85: 3110-3114 (1988); Garin-Chesa et al., Proc. Natl. Acad. USA 87: 7235-7239 (1990); Rettig et al., Cancer Res. 53: 3327-3335 (1993). Each of these four papers is incorporated by reference in its entirety.

Immunohistochemical studies such as those cited supra have shown that $FAP\alpha$ is transiently expressed in certain normal fetal mesenchymal tissues but that normal adult tissues are generally $FAP\alpha^{-}$. Similarly, malignant epithelial, neural and hematopoietic cells are generally $FAP\alpha^{-}$. However, most of the common types of epithelial cancers, including >90% of breast, lung, skin, pancreas, and colorectal carcinomas, contain abundant FAPa reactive stromal fibroblasts. Chesa et al., Proc. Natl. Acad. Sci. USA 87: 7235-7239 (1990). FAPα* tumor stromal fibroblasts almost invariably accompany tumor blood vessels, forming a distinct cellular compartment interposed between the tumor capillary endothelium and the basal aspect of malignant epithelial cell clusters. While $FAP\alpha^+$ stromal fibroblasts are found in both primary and metastatic carcinomas, benign and premalignant epithelial lesions, such as fibroadenomas of the breast and colorectal adenomas only rarely contain FAPa* stromal cells. In contrast

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to the stroma-specific localization of FAP α in epithelial neoplasms, $FAP\alpha$ is expressed in the malignant cells of a large proportion of bone and soft tissue sarcomas. (Rettig et al., Proc. Natl. Acad. Sci. USA 85: 3110-3114 (1988)). $FAPa^*$ fibroblasts have been detected in the granulation tissue of healing wounds (Garin-Chesa et al., supra). Based on the restricted distribution pattern of FAP α in normal tissues and its uniform expression in the supporting stroma of many epithelial cancers, clinical trials with 131I-labeled mAb F19 have been initiated in patients with metastatic colon cancer (Welt et al., Proc. Am. Assoc. Cancer Res. 33: 319 (1992); Welt et al. J. Clin. Oncol. 12: 1561-1571 (1994)) to explore the concept of "tumor stromal targeting" for immunodetection and immunotherapy of epithelial cancers.

Rettig et al., Int. J. Cancer 58: 385-392 (1994), incorporated by reference, discusses the FAP α molecule and its features. Rettig et al postulate that FAP α is found in high molecular weight complexes in excess of 400 kilodaltons, but do not discuss the possibility of dimeric molecules, nor does the paper elaborate on the specific enzymatic properties of the molecule.

The induction of $FAPa^+$ fibroblasts at times and sites of tissue remodeling during fetal development, tissue repair, and carcinogenesis is consistent with a fundamental role for this molecule in normal fibroblast physiology. Thus, it is of interest and value to isolate and to clone nucleic acid molecules which code for this molecule. This is one aspect of the invention, which is described in detail together with other features of the invention, in the disclosure which follows. Further aspects of the invention include the dimeric FAPa molecules, and the exploitation of the properties of these molecules. These features are also elaborated upon hereafter.

BRIEF DESCRIPTION OF THE FIGURES

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Figure 1 compares the deduced amino acid sequence for $FAP\alpha$, and the known sequence of CD26. The alignment has been optimized.

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Figures 2A-2H, inclusive, display immunohistochemical detection of FAP α and CD26 in various tissues. In figures 2A and 2B, breast cancer is studied, for FAP α (figure 2A), and CD26 (figure 2B). In figures 2C and 2D, malignant fibrous histiocytoma is studied, for FAP α (figure 2C), and CD26 (figure 2D). Dermal scar tissue is examined in figures 2E (FAP α), and 2F (CD26). Renal cell carcinoma is studied in figure 2G (FAP α), and 2H (CD26).

Figure 3 presents some of the data generated in experiments which showed that $FAP\alpha$ had extracellular matrix (ECM) protein degrading activity. When zymographic detection of gelatin degrading extracts of 293-FAP was carried out, the active substance was found to have a molecular weight of about 170 kD, via SDS-PAGE, using unboiled samples to preserve enzyme activity.

DETAILED DESCRIPTION OF PREFERRED EMBODIMENTS Example 1

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Fibroblast cell line WI-38 had been observed, previously, to react with mAb F19 (Rettig et al., Canc. Res. 46: 6406-6412 (1986); Rettig et al., Proc. Natl. Acad. USA 85: 3110-3114 (1988); Garin-Chesa et al., Proc. Natl. Acad. Sci. USA 87: 7235-7239 (1990); Rettig et al., Canc. Res. 53: 3327-3335 (1993)). It was used in the experiments which follow.

A cDNA library was prepared from WI-38, using well known available materials. commercially techniques and Specifically, the library was constructed in expression vector pCDNAI, using the Fast Track mRNA isolation kit, and Librarian Once the library was prepared, the cDNA phagemid system. vectors were electroporated into cell line E. coli MC 1061/P3. The pCDNAI expression vector contains an antibiotic resistance gene, so the E. coli were selected via antibiotic resistance. The colonies which were resistant were then used in further experiments. The plasmid DNA from the colonies was obtained via alkaline lysis and purification on CsCl2, in accordance with Sambrook et al, Molecular Cloning: A Laboratory Manual (Cold Spring Harbor Lab, Cold Spring Harbor, N.Y. 2d Ed. The technique is well known to the art, but is

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incorporated by reference herein.

Once the plasmid DNA was isolated, it was used to transfect COS-1 cells, which were then cultured for forty-eight hours, after which these were tested with antibody coated dishes. The mAbs used included F19, as described by Rettig et al., (1986), supra, which is incorporated by reference in its entirety. As COS-1 cells are normally $FAP\alpha^-$, any positive results indicated the presence of the coding sequence. The immunoselection protocol was that of Aruffo et al., Proc. Natl. Acad. Sci USA 84: 3365-3369 (1987), incorporated by reference herein.

Plasmid DNA from positive clones was recovered, in accordance with Hirt, J. Mol. Biol. 26: 365-369 (1967), reintroduced into <u>E. coli</u> MC 1061/P3, and reselected in COS-1 cells.

The protocol presented herein was followed for four rounds. After this, the plasmid DNA of 50 isolated bacterial colonies was purified, using the Qiagen plasmid kit. Of the colonies, 27 clones were found to contain identical 2.8 kb inserts, as determined by EcoRI restriction enzyme mapping. Several of these were found to contain FAPα-specific cDNA as determined by transient expression in COS-1 cells and direct immunofluorescence staining with mAb F19. One of these clones, i.e., "pFAP.38" was selected for further study, as elaborated upon infra.

Example 2

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Once pFAP.38 had been identified, it was tested together with a vector coding for known cell surface marker CD26 ("pCD26"), as well as with control vector pCDNA I.

In these experiments, COS-1 cells were transfected with one of pFAP.38, pCD26, or pCDNAI. After forty-eight hours, the transfectants were tested, using the well known MHA rosetting assay for cell surface antigen expression. In these experiments, mAb F19, which is FAP α specific, was used, together with mAb EF-1, which is CD26 specific. Also used were four other FAP α specific mAbs, i.e., FB23, FB52, FB58 and C48. Also tested were two cancer cell lines, which are known

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to react with mAb F19 (SW872 liposarcoma), or EF-1 (SK-OV6 ovarian cancer). The results are set forth in Table 1, which follows.

Table 1. Cell surface expression of multiple FAP α epitopes and CD26 in human cells and COS-1 cell transfectants

	Cell s	surface a	antigen	express	ion	
Target cell	F19	FB23	FB52	FB58	C48	EF-
Human cells						
SW872 liposarcom	a >95%	>95%	>95%	>95%	>95%	-
SK-OV6 ovarian cancer	-	-	-	-	-	>95
COS-1 transfecta	<u>nts</u>					
COS·pCDNAI control	-	<u>.</u>	-	-	-	-
COS·pFA P 38	40%	30%	40%	20%	20%	-
COS · pCD26	_	-	-	-	-	40

Example 3

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Immunoprecipitation studies were then carried out to identify the antigen being targeted by the antibodies.

Cells were metabolically labelled with Trans ³⁸S-label, (ICN), extracted with lysis buffer (0.01 M Tris-HCl/0.15 M NaCl/0.01 M MgCl₂/0.5% Nonidet P-40/aprotinin (20 ug/ml)/2 mM phenylmethyl-sulfonyl fluoride), and then immunoprecipitated. The protocols used are all well known, as will be seen by reference to Rettig et al., Canc. Res. 53: 3327-3335 (1993); and Fellinger et al., Canc. Res. 51: 336-340 (1991), the disclosures of which are all incorporated by reference in

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Precipitating mAbs were negative control their entirety. mouse Ig, mAb F19, or EF-1. Control tests were carried out with mock transfected COS-1 cells. immunoprecipitation, the immunoprecipitates were boiled in extraction buffer and separated by NaDOdSO./PAGE, reducing conditions. In some experiments, an additional test carried out determine to whether or immunoprecipitated material was glycosylated. In these experiments, cell extracts were fractionated with Con A-SEPHAROSE prior immunoprecipitation. to Following immunoprecipitation, but prior to fractionation NaDodSO4/PAGE, these precipitates were digested with N-Glycanase.

The results showed that, in COS-1 cells, pFAP.38 directs expression of an 88 kd protein species (as determined via SDS-PAGE), which is slightly smaller than the 95 kd FAP α species produced by SW872, or cultured fibroblasts. Digestion with N-Glycanase produced peptides of comparable size (i.e., 74 kd versus 75 kd), showing that the glycosylation of the FAP α protein in COS-1 cells is different than in the human cell lines.

Example 4

Classic Northern blot analysis was then carried out, using the mRNA from FAP α^+ fibroblast cell lines WI-38 and GM 05389, and FAP α^- ovarian cancer cell line SK-OV6. Using the procedures of Sambrook et al., <u>supra</u>, five micrograms of mRNA from each cell line were tested. The probes used were ³²P labelled, and were prepared from a 2.3 kb ECO I fragment of pFAP.38, a 2.4 kb Hind III fragment of CD26, and a 1.8 kb BamHI fragment of γ -actin cDNA. These fragments had been purified from 1% agarose gels.

The extracts of FAP $lpha^*$ fibroblast strains showed a 2.8 kb FAP mRNA species, but extracts of SK-OV6 do not. A γ -actin mRNA species (1.8 kb), was observed in all species.

35 Example 5

The cDNA identified as coding for FAPa was subjected to more detailed analysis, starting with sequencing. The classic

Sanger methodology, as set forth in Proc. Natl. Acad. Sci. USA 74: 5463-5467 (1977), was used to sequence both strands of the cDNA. Once this was secured, an amino acid sequence was deduced therefrom. This information is presented in SEQ ID NO: 1. The sequence was then compared to the known amino acid sequence of CD26 (Morimoto et al., J. Immunol. 143: 3430-3437 (1989)). Figure 1 presents the comparison, using optimized sequence alignment. Any gaps in the comparison are indicated by asterisks, while identical amino acids are shown by dashes in the CD26 sequence. A hydrophobic, putative transmembrane sequence is double underlined, while potential N-glycosylation sites are single underlined.

The sequence analysis shows a 2812 base pair insert, wherein 2277 base pairs constitute the open reading frame. This ORF extends from start codon ATG at nucleotide 209, to stop codon TAA at 2486.

The deduced polypeptide is 760 amino acids long, and has a molecular weight of 87,832. In contrast, N-Glycanase digested, immunopurified FAP α was reported to have an estimated M_r of 75,000 on NaDodSO₄/PAGE (Rettig et al., Canc. Res. 53: 3327-3335 (1993)). A GenBank data base search was carried out. The most closely related genes found were those encoding dipeptidyl peptidase IV homologues (DPPIV; EC 3.4.14.5), with human DPPIV (also known as T-cell activation antigen CD26), showing 61% nucleotide sequence identity, and 48% amino acid sequence identity.

The second set of related genes are human, rat, and bovine homologues of DPPX, a gene of unknown function widely expressed in brain and other normal tissues. The predicted human DPPX gene product shows about 30% amino acid sequence identity with FAP α and CD26. The FAP α molecule exhibits structural features typical of type II integral membrane proteins, including a large COOH-terminal extracellular domain, a hydrophobic transmembrane segment, and a short cytoplasmic tail. The putative extracellular domain contains five potential N-glycosylation sites, eleven cysteine residues (eight of which are conserved between FAP α and CD26), and

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three segments corresponding to highly conserved catalytic domains characteristic of serine proteases, such as DPPIV. These conserved sequences are presented in Table 2, which follows. Comparisons to DPPIV and DPPX were made via Morimoto et al., supra; Wada et al., Proc. Natl. Acad. Sci. USA 89: 197-201 (1992); Yokotani et al., Human Mol. Genet. 2: 1037-1039 (1993).

Example 6

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An additional set of experiments were carried out to determine whether FAP α related sequences are present in non-human species. To do so, human, mouse, and Chinese hamster genomic DNA was digested using restriction enzymes, and tested, via Southern blotting, using the 2.3 kb fragment, labelled with ^{32}P , describes supra. Hybridization was carried out using stringent washing conditions (0.1 x SSC, 0.1% NaDodSO₄, 68°C). Cross-hybridization was readily observed with both the mouse and hamster DNA, suggesting the existence of highly conserved FAP α homologues. In control experiments using the CD26 cDNA fragment described supra, no evidence of cross hybridization was observed.

Example 7

The CD26 molecule shares a number of biochemical and serological properties with FAPB, which is a previously described, FAPa associated molecule having a molecular weight of 105 kd, and is found on cultured fibroblasts melanocytes (Rettig et al., Canc. Res. 53: 3327-3335 (1993)). Cotransfection experiments were carried out to determine whether FAPB is a CD26 gene product. To test this, the same protocols were used which were used for transfection with pFAP.38 or pCD26, as described supra, but using the two vectors. The results presented supra showed cotransfection efficiency was about 40% for each vector, so about 10-20% of cell should be cotransfected.

Following cotransfection, the COS-1 cells were Trans ³⁵S-labeled, as described supra, then lysed, also as described supra.

The resulting cell extracts were separated on Con A

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SEPHAROSE, and the antigen (FAP α and/or CD26) were recovered in the Con A-bound fraction. The bound fraction was eluted with 0.25 M α -D-mannopyranoside. Immunoprecipitation was then carried out, as described supra, and the precipitates were separated on NaDodSO₄/PAGE, also as discussed supra.

Those cells transfected only with pFAP.38 produced FAP α , but not FAP β (determined from mAb F19 immunoprecipitates). They also produce no CD26 antigen (tested with EF-1). Those cells transfected with pCD26 alone produce CD26 but no FAP α . Cotransfectants produce CD26 and FAP α /FAP β heteromers, as determined in the mAb F19 precipitates. This result provides direct evidence that FAP β is a CD26 gene product.

Example 8

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It has been observed previously that some cultured human cell types coexpress FAPa and CD26, and show FAPa/CD26 heteromer formation. In vivo distribution patterns of FAPa in previous determined however, as and CD26. immunohistochemical studies, appeared to be non-overlapping. (See Rettig et al., Proc. Natl. Acad. Sci. USA 85: 3110-3114 (1988); Garin-Chesa et al., Proc. Natl. Acad. Sci. USA 87: 7235-7329 (1990); Rettig et al., Canc. Res. 53: 3327-3335 (1993); Stein et al., in Knapp et al., eds. Leukocyte typing IV-white cell differentiation antigens, pp 412-415 (Oxford University Press, N.Y. 1989), pp. 412-415; Möbious et al., J. Exp. Immunol. 74: 431-437 (1988)). In view of the potential significance of FAPa/CD26 coassociation, tissue distribution was reexamined, via side by side immunohistochemical staining of normal tissues and lesional tissues known to contain $FAPa^*$ fibroblasts or FAPa malignant cells.

To test the samples, they were embedded in OCT compound, frozen in isopentane precooled in liquid nitrogen, and stored at -70°C until used. Five micrometer thick sections were cut, mounted on poly-L-lysine coated slides, air dried, and fixed in cold acetone (4°C, for 10 minutes). The sections were then tested with mAbs (10-20 ug/ml), using the well known avidin-biotin immmuno-peroxidase method, as described by, e.g., Garin-Chesa et al., J. Histochem. Cytochem. 37: 1767-

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1776 (1989); Garin-Chesa et al., Proc. Natl. Acad. Sci. USA 87: 7235-7239 (1990); Rettig et al., Canc. Res. 53: 3327-3335 (1993); Garin-Chesa et al., Am. J. Pathol. 142: 557-567.

The results are shown in figure 2. Breast, colorectal, pancreas and lung carcinomas showed strong expression of FAP α and no CD26 was found (see figures 2A and 2B). Five FAP α sarcomas, including malignant fibrous histiocytoma (figures 2C and 2D), were tested, and there was no expression of CD26. Examination of reactive fibroblasts of healing dermal wounds (figures 2E, 2F), showed abundant expression of both FAP α and CD26. The three renal carcinomas tested (figures 2G, 2H), showed expression of CD26 in malignant epithelium. FAP α was absent from malignant epithelial cells, and showed low expression in the stroma of these carcinomas.

Example 9

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A mammalian cell line, transfected with a $\textsc{FAP}\alpha$ encoding cDNA, was prepared.

Human embryonic kidney cell line 293 is well known and widely available from, e.g., the American Type Culture Collection.

Samples of 293 were maintained, in an incubator, at 37°C, in an atmosphere of 95% air, and 5% CO₂. The cells were cultured in a 50:50 mixture of Dulbecco's modified minimal essential medium and Ham's F12 medium, augmented with 10% fetal bovine serum, penicillin and streptomycin. Following the procedures described by Ustar et al., Eur. Mol. Biol. J. 1991, and/or Park et al., J. Biol. Chem. 169: 25646-25654 (1994), both of which are incorporated by reference, cDNA for FAPa (i.e., SEQ ID NO: 1), was transfected into the 293 cells. Details of the cDNA vector are provided, supra (pFAP.38). Transfectants were selected for resistance to antibiotics (200 ug/ml Geneticin), and were then maintained in selection medium, containing Geneticin.

Individual colonies of resistant cells were picked, grown to confluence in 6 well tissue culture plates, and were tested for FAP α expression in an immunofluorescence assay (IFA), using FAP α specific monoclonal antibody F19 as described

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supra.

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Those colonies which expressed $FAP\alpha$ were expanded, and monitored by indirect IFA and cytofluorometric analysis, also as set forth, supra.

The IFAs were positive for the transfectants, referred to hereafter as cell line 293-FAP, but were negative for parental line 293.

Example 10

In order to confirm that recombinant FAPa was, in fact, being produced, a series of immunoprecipitation experiments were carried out. These followed the methods of Park, et al., supra, and Rettig et al., Canc. Res. 53: 3327-3335 (1993), both of which are incorporated by reference. Essentially, both of which are incorporated by reference. Essentially, methionine labelled cell extracts were combined with monoclonal antibody F19, in the manner described supra. Precipitates were then boiled in extraction buffer and run on SDS-PAGE gels, using, as a negative control, mouse IgG1. Both cell line 293-FAP, and non transfected line 293 were tested. The results indicated clearly, that recombinant FAPa was produced by the transfected cell line 293-FAP. This was determined by immunoprecipitation analyses, using FAPa specific monoclonal antibody F19.

Example 11

The ability to produce recombinant FAP α permitted further study of the molecule's properties. Specifically, given the structural features outlined in the prior examples, experiments were designed to determine if FAP α possesses enzymatic activities. The experiments were designed to test whether or not FAP α had extracellular matrix (ECM) protein degrading activity.

Extracts of 293-FAP cells were prepared, using an extraction buffer (0.15M NaCl, 0.05M Tris-HCl, pH 7.4, 10 mM MgCl₂, 1 percent Triton X-114), were cleared by centrifugation (4,000xg, 10 minutes at 4°C), and phase partitioned at 37°C for 10-20 minutes. This was followed by further centrifugation (4000xg, 20 minutes at 20-25°C). Detergent phases were diluted with buffer (0.15 M NaCl, 0.05 M Tris-HCl

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pH 7.4, 5 mM CaCl₂, 5 mM MgCl₂, 0.75% Empigen BB), and separated on concanavalin A-Sepharose following Rettig et al., supra. Any concanavalin A bound fractions were eluted with 0.25M methyl-α-D-mannopyranoside in elution buffer 0.15 M NaCl₂,0.05 M Tris-HCl₂, pH 7.4, 5mM CaCl₂, 5 mM MgCl₂, 0.1% Triton X-100), mixed with zymography sample buffer (0.25 M Tris-HCl₂, pH 6.8, 8% SDS, 40% glycerol, 0.01% bromophenol blue), at a 3:1 ratio, and used for further analysis.

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Aliquots of sample were loaded onto polyacrylamide gels containing 0.1% of either of gelatin or Electrophoresis was then carried out in a Biorad Mini-Protein II system, at 20 mA constant current for 1.5 - 2 hours, until the bromophenol blue due fronts of samples had reached the lower end of the gel. The gel was removed and incubated for one hour at 20-25°C in a 2.5% aqueous solution of Triton X-100 on a rotary shaker. The Triton X-100 solution was decanted, and replaced with enzyme buffer (0.05M Tris-HCl, pH 7.5, 0.2M NaCl, 5 mM CaCl₂, 5 mM MgCl₂, 0.02% Brij 35). The gel was then incubated at 37°C or 41°C, followed by staining or destaining at room temperature. Gels were stained with 0.5% of Coomassie Brilliant Blue G-250 in an aqueous solution of 30% methanol and 10% acetic acid for 15, 30, and 60 minutes, respectively. Subsequently, gels were incubated for 15 minutes in an aqueous solution of 30% CH,OH and 5% glycerol, followed by drying between sheets of cellophane.

Gelatinase activity was evaluated in accordance with Kleiner et al., Anal. Biochem. 218: 325-329 (1994), incorporated by reference in its entirety. This is a routine assay used to determine whether or not a protease capable of digesting gelatin is present. Labelled molecular weight standard were run on the same gels, under reducing conditions, for molecular weight determinations.

Proteolytic activity for defined amino acid sequence motifs were tested, using a well known membrane overlay assay. See Smith et al, Histochem. J. 24(9): 637-647 (1992), incorporated by reference. Substrates were Ala-Pro-7-amino-4-trifluoromethyl coumarin, Gly-Pro-7-amino-4-trifluoromethyl

coumarin, and Lys-Pro-7-amino-4-trifluromethyl coumarin.

The results of these experiments are depicted, in part, in figure 3. This figure shows zymographic detection of gelatin degrading activity, in the cell extracts. See Kleiner et al., supra. A protein species of approximately 170 kilodaltons, as determined by SDS-PAGE, was observed to have gelatin degrading activity. This species, which was found in the 293-FAP cell line, but not in untransfected 293 cells, is thus identified as FAPa. The molecular weight is consistent with a dimer, i.e., a dimeric FAPa molecule.

The proteolytic activity described herein where gelatin is the substrate, was not observed when casein was the substrate.

Example 12

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Further studies were then undertaken in order to characterize the 170 kD FAPa dimer further. Specifically, the experiments described in example 11 were repeated, except that 5% of 2-mercaptoethanol or 5 um iodoacetamide was added to the extracts prior to SDS-PAGE, or ethylenediamine N,N,N',N'-tetraacetic acid (10 mM) was added to the incubation buffer used for gelatin zymography. None of these treatments abolished the enzymatic activity. In contrast, heating at 100°C for five minutes prior to SDS-polyacrylamide gel electrophoresis abolished the gelatin-degrading activity.

Further work, using a membrane overlay assay, described by, e.g., Smith et al., Histochem J. 24(9): 643-647 (1992), incorporated by reference, revealed that the FAPα dimers were able to cleave all of the Ala-Pro, Gly-Pro, and Lys-Pro dipeptides tested.

In further experiments, a fusion protein was produced which comprised the extracellular domains of both FAP α and murine CD8 proteins. This chimeric protein was produced in a baculovirus system in insect cells. The chimeric protein exhibited the same enzymatic activity as FAP α , using the model discussed supra.

Example 13

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Two quantitative assays for $FAP\alpha$ enzyme activity were developed using Ala-Pro-7-amino-4-trifluoromethyl coumarin (Ala-Pro-AFC) as the substrate. In the first assay format, membrane extracts of FAP α -expressing cells were mixed with a 5-10 fold volume of reaction buffer (100mM NaCl, 100mM Tris pH 7.8), and added to an equal volume of 0.5mM Ala-Pro-AFC in reaction buffer followed by an incubation for one hour at 37°C. Release of free AFC was then measured in a fluorimeter using a 395nm excitation / 530nm emission filter set. membrane extracts analyzed in this assay format were derived from either 293-FAP α cells (293 cells stably transfected with vector FAP.38 described supra) or HT1080-FAPα cells (HT1080 cells stably transfected with vector FAP.38). Negative control experiments assessing FAPa-specific activities were carried out with membrane extracts prepared respective parental 293 or HT1080 cell lines. In the second assay, FAP α was isolated from 293-FAP α or HT1080-FAP α membrane extracts via an antibody specific for FAP α . Ninety-six well ELISA plates were coated overnight at 4°C with lug/ml F19 monoclonal antibody in phosphate-buffered saline (PBS). the case of CD8-FAP α discussed infra plates were coated with F19 antibody as above or with lug/ml rat anti-mouse CD8 overnight at 4°C. Wells were then washed with wash buffer (PBS, 0.1% Tween 20). Excess binding sites were blocked with blocking buffer (5% bovine serum albumin in PBS) for 1 hour at room temperature. Blocking buffer was removed; membrane extracts of 293-FAP α expressing cells or control cells were added and incubated for 1 hour at room temperature. unbound material was removed, wells were washed with wash buffer, and FAPa activity was assayed using 100 ul Ala-Pro-AFC (0.5 mM Ala-Pro-AFC in reaction buffer) for one hour at 37°C. Release of free AFC was measured as above. Binding of mab F19 to FAPa did not measurably affect its enzymatic activity.

Example 14

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Using assays for FAPa enzyme activity, described supra an inhibitor of FAP α enzymatic activity has been identified. This inhibitor is (S)-Valylpyrrolidine-2(R)-boronic acid (Snow

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et al., J. Am. Chem Soc.(1994) 116:10860-10869), referred to here as ValboroPro. ValboroPro inhibits cleavage of Ala-Pro-AFC by FAP α with an IC $_{50}$ of 0.11 uM. ValboroPro also inhibits the gelatinolytic activity of FAP α at a concentration of 100 uM. The specificity of ValboroPro for FAP α was demonstrated in tests with an unrelated serine protease, trypsin. No inhibition of bovine trypsin by ValboroPro (up to 100 uM) was observed when assayed with carbobenzoxy-L-valinyl-glycinyl-L-arginyl-4-nitranilide acetate as substrate.

Example 15

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The identification of specific, structural requirements the enzymatic activities of $FAP\alpha$ facilitates the development of molecules which can bind to and/or inhibit To examine whether the serine residue at position 624 of the predicted amino acid sequence of FAPa polypeptide is critical for its enzymatic function, site-directed mutagenesis according to Zoller, et al DNA 3:479-488 (1984) was performed using standard polymerase chain reaction methods. codon coding for serine 624 in the FAP α cDNA was replaced with GCG, resulting in alanine at this position. The altered DNA was reintroduced into the FAP.38 vector and transfected into 293 cells as described supra. Geneticin-resistant colonies were selected and examined by indirect IFA for FAPlphaexpression using mAb F19 as well as other FAP α specific antibodies described by Rettig, et al., J. Cancer 58:385-392 (1994) as set forth, supra. No differences in binding of the anti-FAPa antibodies to the mutant FAPa expressing cells were observed as compared to wild type FAPa transfected cells. The presence of the mutation was confirmed through amplification of genomic DNA and restriction enzyme digestion performed with several clones of transfected cells. To assess the enzymatic activity of mutant FAPa, the following tests were performed. Membrane extracts were prepared from three positive clones and equal amounts of FAPaprotein (as determined in a double-determinate ELISA assay using two anti-FAPa antibodies that recognize distinct FAPa epitopes) were examined in the gelatinolytic and Ala-Pro-AFC capture assays.

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Both the gelatinolytic activity and the activity in the capture assay of isolated mutant $FAP\alpha$ were reduced to undetectable levels compared to wild type $FAP\alpha$, confirming the role of the canonical serine in the catalytic triad for both observed enzymatic activities.

Example 16

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A fusion protein was generated to obtain secreted, water-FAPα soluble enzyme. In this fusion protein, the extracellular domain of CD8, consisting of the first 189 amino acids of murine CD8, was linked to the extracellular domain of FAPa (amino acids 27 to 760), as described by Lane et al., J. Exp. Med. 177:1209 (1993)using standard polymerase chain reaction protocols and inserted in commercially available pVL1393 vector. Transfection of Sf9 cells with this vector and amplification of the resulting recombinant baculovirus were performed as described (Baculovirus Expression Vectors. O'Reilly, Miller, and Luckow, Oxford University Press, 1994). fusion protein was isolated in a two step The CD8-FAP purification from the spent medium of High Five™ cells infected with CD8-FAPa baculovirus for four days. Cells and virus were removed by ultracentrifugation, the supernatant was passed through а column containing Heparin-Sepharose (Pharmacia) and eluted stepwise with 0.6, 1.0, and 2.0 M NaCl in 10mM phosphate, pH 7. Active fractions from the 1.0 and 2.0 M eluates were pooled and concentrated using an YM-10 filter and 26/60 Superdex-200 gel filtration column. Activity was observed in a high molecular weight peak which, when subjected to N-terminal gas phase sequencing, was confirmed to be CD8-FAPa. In gelatinolytic assays, activity greater than 200kD in the gelatinolytic assay was detected when purifieed CD8-FAPa was tested, consistent with the higher predicted molecular weight of the fusion protein.

Example 17

The presence of structural and functional homologues in non-human species has been ascertained. For example, the cDNA for mouse FAP α has been cloned and characterized. Examination of the predicted amino acid sequence of the homologous mouse

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FAP α cDNA sequence (EMBL accession number Y10007) reveals a high degree of conservation of FAP α across species. The two proteins are 89% identical and the catalytic triad is conserved between human FAP α and mouse FAP α . The high degree of conservation and similar tissue expression suggests that FAP α from nonhuman sources may be functionally equivalent to human FAP α . This conclusion is confirmed by the finding that a CD8-murine FAP α fusion protein similar in design to CD8-human FAP α also demonstrates the expected dipeptidylpeptidase enzymatic activity using Ala-Pro-AFC as substrate.

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The foregoing examples describe an isolated nucleic acid molecule which codes for fibroblast activating protein alpha ("FAPa"), as well as dimeric forms of the molecule, and uses thereof. The expression product of the sequence in COS-1 is a protein which, on SDS-PAGE of boiled samples, shows a molecular weight of about 88 kd. Deduced amino acid sequence, as provided in SEQ ID NO: 1, for one form of the molecule, yields a molecular weight of about 88 kd.

It should be noted that there is an apparent discrepancy in molecular weight in that the COS-1 isolate is glycosylated, while molecular weight from deduced amino acid sequences does not account for glycosylation. Membrane proteins are known to exhibit aberrant migration in gel systems, however, which may explain the difference observed here.

Also a part of the invention are chimeric and fusion proteins, which comprise a portion of FAP α which contain the molecule's catalytic domain, and additional, non FAP α components. The FAP α catalytic domain <u>per se</u> is also a part of the invention.

It is to be understood that, as described, FAP α may be glycosylated, with the type and amount of glycosylation varying, depending upon the type of cell expressing the molecule. The experiment described herein shows this. This is also true for the dimeric form of the molecule, first described herein, having a molecular weight of about 170 kilodaltons as determined by SDS-PAGE of unboiled samples.

The invention also comprehends the production of

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expression vectors useful in producing the FAP α molecule. In their broadest aspect, these vectors comprise the entire FAP α coding sequence or portions thereof, operably linked to a promoter. Additional elements may be a part of the expression vector, such as protein domains fused to the FAP α protein or protein portions ("fusion protein") genes which confer antibiotic resistance, amplifiable genes, and so forth.

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The coding sequences and vectors may also be used to prepare cell lines, wherein the coding sequence or expression vector is used to transfect or to transform a recipient host. The type of cell used may be prokaryotic, such as <u>E. coli</u>, or eukaryotes, such as yeast, CHO, COS, or other cell types.

The identification of nucleic acid molecules such as that set forth in SEQ ID NO: 1 also enables the artisan to identify and to isolate those nucleic acid molecules which hybridize to it under stringent conditions. "Stringent condition" as used herein, refers to those parameters set forth supra, whereby both murine and hamster sequences were also identified. It will be recognized by the skilled artisan that these conditions afford a degree of stringency which can be achieved using parameters which vary from those recited. Such variance is apprehended by the expression "stringent conditions".

The ability of nucleic acid molecules to hybridize to complementary molecules also enables the artisan to identify cells which express $FAP\alpha$, via the use of a nucleic acid hybridization assay. One may use the sequences described in the invention to hybridize to complementary sequences, and thus identify them. In this way, one can target mRNA, e.g., which is present in any cell expressing the $FAP\alpha$ molecule.

It is of course understood that the nucleic acid molecules of the invention are also useful in the production of recombinant FAPa, in both monomeric and dimeric form. The examples clearly show that host cells are capable of assembling the dimeric forms. The recombinant protein may be used, e.g., as a source of an immunogen for generation of antibodies akin to known mAb F19, and with the same uses. Similarly, the recombinant protein, and/or cells which express

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the molecule on their surface, may be used in assays to determine antagonists, agonists, or other molecules which interact with molecules having FAP α activity. Such substances may be, but are not necessarily limited to, substrates, inhibiting molecules, antibodies, and so forth. The molecules having FAP α activity may be, e.g., the monomeric or dimeric forms of FAPa, derivatives containing the catalytic domain, and so forth. The molecule having $FAP\alpha$ activity may be pure, or in the form of a cell extract, such as a transformed or transfected cell, which has received an FAP α active gene. Both prokaryotes and eukaryotes may be used. feature of the invention should be considered in light of the observed structural resemblances to membrane bound enzymes. This type of molecule is associated with certain properties which need not be described in detail here. It will suffice to say that inhibition or potentiation of these properties as associated with $FAP\alpha$ is a feature of this invention. example, one may identify substrates or the substrate for FAP α molecules, via the use of recombinant cells or recombinant FAPα per se. The substrates can be modified to improve their effect, to lessen their effect, or simply to label them with detectable signals so that they can be used, e.g., to identify Study of the interaction of cells which express FAPa. substrate and FAP α , as well as that between FAP α and any molecule whatsoever, can be used to develop and/or to identify agonists and antagonists of the FAP α molecule.

Also a feature of the invention are isolated, dimeric $FAP\alpha$

molecules which have a molecular weight of about 170 kilodaltons as determined by SDS-PAGE, their use as an enzymatic cleaving agent, and other uses as are described herein. Enzymatically active forms of FAP α may also be produced as recombinant fusion proteins, such as soluble fusion proteins comprising the catalytic domain of FAP α and other protein domains with suitable biochemical properties, including secretory signals, protease cleavage sites, tags for purification, and other elements known to the artisan.

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Exemplary are CD8 peptide sequences, such as are described <u>supra</u>. The fact that FAP α has particular properties, as described herein, permits the identification of the molecule on cells expressing them. In turn, because the FAP α molecule is associated with tumors and tumor stromal cells, targeting of FAP α with therapeutic agents serves as a way to treat cancerous or precancerous condition, by administering sufficient therapeutic agent to alleviate cancer load.

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The experiments showing the proteolytic properties of FAPa lead to yet a further aspect of the invention. well known that proteases which degrade extracellular matrix, or "ECM" proteins have an important role on certain aspects of tumor growth, including their effect on tumor cell invasion, tumor blood vessel formation (i.e., neoangiogenesis), and tumor metastasis. Collagens are of special interest vis-a-vis the substrates of proteases, as the collagens are an important part of the ECM. The fact that FAPa digests ECM suggests a therapeutic role for inhibitors of the molecule. "Inhibitors", as used herein, refers to molecules which interfere with FAPa enzyme function. Specifically excluded from such inhibitors is the monoclonal antibody F19. This mAb is known to bind to but not inhibit the enzyme function of FAPa, and hence it is not an inhibitor. The art is quite well versed with respect to monoclonal antibodies which both bind to and inhibit enzymes. Further examples of such inhibitors would include, e.g., substrate derivatives, such as modified collagen molecules, which interfere with the active site or sites of the FAPa molecule. Other suitable inhibitors will be apparent to the skilled artisan, and need not be listed here. addition, the recombinant FAPa proteins transfected cell lines described supra can be employed in an enzymatic screening assay, using the substrate described supra or other suitable substrates, to identify inhibitors from any The identification of substances which compound library. interact with FAPa active molecules thereby leads therapeutic treatment of conditions where a subject exhibit abnormal FAPa activity. Specifically, an amount of an

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appropriate substance, be it an inhibitor (e.g, a collagen derivative, S-Valyl-pyrrolidine-2(R)-boronic acid), an agonist or an antogonist is administered to a subject in an amount sufficient to normalize FAP α activity.

Other aspects of the invention will be clear to the skilled artisan, and need not be set forth here.

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The terms and expressions which have been employed are used as terms of description and not of limitation, and there is no intention in the use of such terms and expressions of excluding any equivalents of the features shown and described or portions thereof, it being recognized that various modifications are possible within the scope of the invention.

PCT/US97/04215 WO 97/34927

(1) GENERAL INFORMATION:
(i) APPLICANTS: Zimmermann, Rainer; Park, John E.;
Rettig, Wolfgang; Old, Lloyd J.
(ii) TITLE OF INVENTION: ISOLATED DIMERIC FIBROBLAST ACTIVATION
PROTEIN ALPHA, AND USES THEREOF
(iii) NUMBER OF SEQUENCES: 2
(iv) CORRESPONDENCE ADDRESS:
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(B) STREET: 805 Third Avenue
(C) CITY: New York City
(D) STATE: New York
(E) COUNTRY: USA
(F) ZIP: 10022
(v) COMPUTER READABLE FORM:
(A) MEDIUM TYPE: Diskette, 3.5 inch, 2.0 MB storage
(B) COMPUTER: IBM PS/2
(C) OPERATING SYSTEM: PC-DOS
(D) SOFTWARE: Wordperfect
(vi) CURRENT APPLICATION DATA:
(A) APPLICATION NUMBER:
(B) FILING DATE:
(C) CLASSIFICATION:
(vii)PRIOR APPLICATION DATA:
(A) APPLICATION NUMBER: 08/619,280
(B) FILING DATE: 18-MARCH-1996
(C) CLASSIFICATION: 435
(vii) PRIOR APPLICATION DATA:
(A) APPLICATION NUMBER: 08/230,491
(B) FILING DATE: 20-APRIL-1994
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(B) REGISTRATION NUMBER: 30,946
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- (2) INFORMATION FOR SEQ ID NO: 1:
 - SEQUENCE CHARACTERISTICS:

(A) LENGTH:

2815 Base pairs

(B) TYPE:

nucleic acid

(C) STRANDEDNESS: double

(D) TOPOLOGY:

linear

(xi) SEQUENCE DESCRIPTION: SEQ. ID NO: 1:

AAGAACGCCC CCAAAATCTG TTTCTAATTT TACAGAAATC TTTTGAAACT TGGCACGGTA 60 10 TTCAAAAGTC CGTGGAAAGA AAAAAACCTT GTCCTGGCTT CAGCTTCCAA CTACAAAGAC 120 AGACTTGGTC CTTTTCAACG GTTTTCACAG ATCCAGTGAC CCACGCTCTG AAGACAGAAT 180 TAGCTAACTT TCAAAAACAT CTGGAAAAAT GAAGACTTGG GTAAAAATCG TATTTGGAGT 240 TGCCACCTCT GCTGTGCTTG CCTTATTGGT GATGTGCATT GTCTTACGCC CTTCAAGAGT 300 TCATAACTCT GAAGAAAATA CAATGAGAGC ACTCACACTG AAGGATATTT TAAATGGAAC 360 15 ATTTTCTTAT AAAACATTTT TTCCAAACTG GATTTCAGGA CAAGAATATC TTCATCAATC 420 TGCAGATAAC AATATAGTAC TTTATAATAT TGAAACAGGA CAATCATATA CCATTTTGAG 480 TAATAGAACC ATGAAAAGTG TGAATGCTTC AAATTACGGC TTATCACCTG ATCGGCAATT 540 TGTATATCTA GAAAGTGATT ATTCAAAGCT TTGGAGATAC TCTTACACAG CAACATATTA 600 CATCTATGAC CTTAGCAATG GAGAATTTGT AAGAGGAAAT GAGCTTCCTC GTCCAATTCA 20 660 GTATTTATGC TGGTCGCCTG TTGGGAGTAA ATTAGCATAT GTCTATCAAA ACAATATCTA 720 TTTGAAACAA AGACCAGGAG ATCCACCTTT TCAAATAACA TTTAATGGAA GAGAAAATAA 780 AATATTTAAT GGAATCCCAG ACTGGGTTTA TGAAGAGGAA ATGCTTCCTA CAAAATATGC 840 TCTCTGGTGG TCTCCTAATG GAAAATTTTT GGCATATGCG GAATTTAATG ATAAGGATAT 900 ACCAGTTATT GCCTATTCCT ATTATGGCGA TGAACAATAT CCTAGAACAA TAAATATTCC 960 25 ATACCCAAAG GCTGGAGCTA AGAATCCCGT TGTTCGGATA TTTATTATCG ATACCACTTA 1020 CCCTGCGTAT GTAGGTCCCC AGGAAGTGCC TGTTCCAGCA ATGATAGCCT CAAGTGATTA 1080 TTATTTCAGT TGGCTCACGT GGGTTACTGA TGAACGAGTA TGTTTGCAGT GGCTAAAAAG 1140 AGTCCAGAAT GTTTCGGTCC TGTCTATATG TGACTTCAGG GAAGACTGGC AGACATGGGA 1200 TTGTCCAAAG ACCCAGGAGC ATATAGAAGA AAGCAGAACT GGATGGGCTG GTGGATTCTT 1260 30 TGTTTCAAGA CCAGTTTTCA GCTATGATGC CATTTCGTAC TACAAAATAT TTAGTGACAA 1320 GGATGGCTAC AAACATATTC ACTATATCAA AGACACTGTG GAAAATGCTA TTCAAATTAC 1380 AAGTGGCAAG TGGGAGGCCA TAAATATATT CAGAGTAACA CAGGATTCAC TGTTTTATTC 1440 TAGCAATGAA TTTGAAGAAT ACCCTGGAAG AAGAAACATC TACAGAATTA GCATTGGAAG 1500 CTATCCTCCA AGCAAGAAGT GTGTTACTTG CCATCTAAGG AAAGAAAGGT GCCAATATTA 1560 35 CACAGCAAGT TTCAGCGACT ACGCCAAGTA CTATGCACTT GTCTGCTACG GCCCAGGCAT 1620 CCCCATTTCC ACCCTTCATG ATGGACGCAC TGATCAAGAA ATTAAAATCC TGGAAGAAAA 1680

CAAGGAATTG GAAAATGCTT TGAAAAATAT CCAGCTGCCT AAAGAGGAAA TTAAGAAACT 1740 TGAAGTAGAT GAAATTACTT TATGGTACAA GATGATTCTT CCTCCTCAAT TTGACAGATC 1800 AAAGAAGTAT CCCTTGCTAA TTCAAGTGTA TGGTGGTCCC TGCAGTCAGA GTGTAAGGTC 1860 TGTATTTGCT GTTAATTGGA TATCTTATCT TGCAAGTAAG GAAGGGATGG TCATTGCCTT 1920 GGTGGATGGT CGAGGAACAG CTTTCCAAGG TGACAAACTC CTCTATGCAG TGTATCGAAA 1980 GCTGGGTGTT TATGAAGTTG AAGACCAGAT TACAGCTGTC AGAAAATTCA TAGAAATGGG 2040 TTTCATTGAT GAAAAAAGAA TAGCCATATG GGGCTGGTCC TATGGAGGAT ACGTTTCATC 2100 ACTGGCCCTT GCATCTGGAA CTGGTCTTTT CAAATGTGGT ATAGCAGTGG CTCCAGTCTC 2160 CAGCTGGGAA TATTACGCGT CTGTCTACAC AGAGAGATTC ATGGGTCTCC CAACAAAGGA 2220 TGATAATCTT GAGCACTATA AGAATTCAAC TGTGATGGCA AGAGCAGAAT ATTTCAGAAA 2280 TGTAGACTAT CTTCTCATCC ACGGAACAGC AGATGATAAT GTGCACTTTC AAAACTCAGC 2340 ACAGATTGCT AAAGCTCTGG TTAATGCACA AGTGGATTTC CAGGCAATGT GGTACTCTGA 2400 CCAGAACCAC GGCTTATCCG GCCTGTCCAC GAACCACTTA TACACCCACA TGACCCACTT 2460 CCTAAAGCAG TGTTTCTCTT TGTCAGACTA AAAACGATGC AGATGCAAGC CTGTATCAGA 2520 ATCTGAAAAC CTTATATAAA CCCCTCAGAC AGTTTGCTTA TTTTATTTTT TATGTTGTAA 2580 AATGCTAGTA TAAACAAACA AATTAATGTT GTTCTAAAGG CTGTTAAAAA AAAGATGAGG 2640 ACTCAGAAGT TCAAGCTAAA TATTGTTTAC ATTTTCTGGT ACTCTGTGAA AGAAGAGAAA 2700 AGGGAGTCAT GCATTTTGCT TTGGACACAG TGTTTTATCA CCTGTTCATT TGAAGAAAAA 2760 TAATAAAGTC AGAAGTTCAA AAAAAAAAAA AAAAAAAAA AAAGCGGCCG CTCGA 2815

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(2) INFORMATION FOR SEQ ID NO: 2:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH:

760 amino acids

(B) TYPE:

amino acid

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION:

SEQ ID NO: 2:

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Met Lys Thr Trp Val Lys Ile Val Phe Gly Val Ala Thr Ser Ala Val

5 10 15

Leu Ala Leu Leu Val Met Cys Ile Val Leu Arg Pro Ser Arg Val His
20 25 30

Asn Ser Glu Glu Asn Thr Met Arg Ala Leu Thr Leu Lys Asp Ile Leu
35 40 45

Asn Gly Thr Phe Ser Tyr Lys Thr Phe Phe Pro Asn Trp Ile Ser Gly

		50					55					60				
	G 1 n 65	G1u	Tyr	Leu	His	G 1 n 70	Ser	Ala	Asp	Asn	Asn 75	Ile	Va 1	Leu	Tyr	Asn 80
5		Glu	Thr	Gly	G1n 85	Ser	Tyr	Thr	I 1e	Leu 90	Ser	Asn	Arg	Thr	Met 95	Lys
5	Ser	Va 1	Asn	Ala 100		Asn	Tyr	Gly	Leu 105	Ser	Pro	Asp	Arg	Gln 110	Phe	Val
	Tyr	Leu	G1u 115	Ser	Asp	Tyr	Ser	Lys 120	Leu	Trp	Arg	Tyr	Ser 125	Tyr	Thr	Ala
10	Thr	Tyr 130	Tyr	I le	Tyr	Asp	Leu 135	Ser	Asn	Gly	Glu	Phe 140	Val	Arg	Gly	Asn
	145					150				Cys	155					160
15					165					I le 170					175	
			•	180		•			185	Asn				190		
			195					200		Glu			205			
20		210					215			Gly		220				
	225					230				Ile	235					240
25					245			•		I le 250					255	
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30		290)				295			Trp		300				
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35		·			325	•				7 Trp)				335	•
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	Ser	Arg		Val	Phe	Ser	Tyr			Ile	Ser	Tyr	Tyr	-	Ile	Phe
			355	_		_		360					365			•
	Ser	Asp 370	Lys	Asp	Gly	Tyr	Lys 375	His	Ile	His	Tyr	I le 380	Lys	Asp	Thr	' Va`
5	Glu	Asn	Ala	I le	Gln	I le	Thr	Ser	Gly	Lys	Trp		Ala	I le	Asn	H
	385					390			_		395					400
			۷al	Thr	Gln	Asp	Ser	Leu	Phe	Tyr			Asn	Glu	Phe	
		_			405	·				410					415	
	G lu	Tyr	Pro	Gly	Arg	Arg	Asn	Ile	Tyr			Ser	I le	Gly		
10				420					425					430		
	Pro	Pro	Ser	Lys	Lys	Cys	Va 1	Thr	Cys	His	Leu	Arg	Lys	Glu	Arq	Cys
			435					440	-				445		J	
	Gln	Tyr	Tyr	Thr	Ala	Ser	Phe	Ser	Asp	Tyr	Ala	Lys	Tyr	Tyr	Ala	Leu
		450					455					460				
15	Va 1	Cys	Tyr	Gly	Pro	Gly	Ile	Pro	I le	Ser	Thr	Leu	His	Asp	Gly	Arg
	465					470					475					480
	Thr	Asp	Gln	Glu	Ile	Lys	I le	Leu	Glu	Glu	Asn	Lys	Glu	Leu	Glu	Asn
					485					490					495	
	Ala	Leu	Lys	Asn	Пe	Gln	Leu	Pro	Lys	Glu	Glu	Пe	Lys	Lys	Leu	G٦u
20				500					505					510		
	Va 1	Asp	Glu	I le	Thr	Leu	Trp	Tyr	Lys	Met	Ile	Leu	Pro	Pro	Gln	Phe
			515					520					525			
	Asp	Arg	Ser	Lys	Lys	Tyr	Pro	Leu	Leu	I le	Gln	Va 1	Tyr	Gly	Gly	Pro
		530					535					540				
25		Ser	61n	Ser	Va 1		Ser	Va 1	Phe	Ala	Va 1	Asn	Trp	I le	Ser	Tyr
	545					550					555					560
	Leu	Ala	Ser	Lys		Gly	Met	Val	I le		Leu	Va 1	Asp	Gly	Arg	Gly
		_			565					570					575	
	Thr	Ala	Phe		Gly	Asp	Lys	Leu		Tyr	Ala	Val	Tyr	_	Lys	Leu
30				580					585					590		
	Gly	Va 1		Glu	Val	Glu	Asp		Ile	Thr	Ala	Val	Arg	Lys	Phe	Ile
			595					600					605			
	Glu		Gly	Phe	Ile	Asp		Lys	Arg	He	Ala		Trp	Gly	Trp	Ser
	_	610		_			615					620				
35		Gly	Gly	Tyr	Va 1		Ser	Leu	Ala	Leu		Ser	Gly	Thr	Gly	
	625		_			630			_		635		_			640
	Phe	Lys	Cys	Gly	Ile	Ala	Va 1	Ala	Pro	Va 1	Ser	Ser	Trp	Glu	Tyr	Tyr

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					645					650					655	
	Ala	Ser	Va 1	Tyr	Thr	Glu	Arg	Phe	Met	Gly	Leu	Pro	Thr	Lys	Asp	Asp
				660					665					670		
	Asn	Leu	Glu	His	Tyr	Lys	Asn	Ser	Thr	Va 1	Met	Ala	Arg	Ala	Glu	Tyr
5			675					680					685			
	Phe	Arg	Asn	Val	Asp	Tyr	Leu	Leu	Ile	His	Gly	Thr	Ala	Asp	Asp	Asn
		690					695					700				
	Va 1	His	Phe	Gln	Asn	Ser	Ala	Gln	I le	Ala	Lys	Ala	Leu	Va 1	Asn	Ala
	705					710					715					720
10	G I n	Val	Asp	Phe	Gln	Ala	Met	Trp	Tyr	Ser	Asp	Gln	Asn	His	Gly	Leu
					725					730					735	
	Ser	Gly	Leu	Ser	Thr	Asn	His	Leu	Tyr	Thr	His	Met	Thr	His	Phe	Leu
				740					745					750		
	Lys	Gln	Cys	Phe	Ser	Leu	Ser	Asp								
15			755					760								

We claim:

- Isolated, dimeric FAPα molecule, having a molecular weight of about 170 kilodaltons as determined by SDS-PAGE, wherein said dimeric FAPα molecule is capable of degrading extracellular matrix proteins.
- 2. The dimeric FAP α molecule of claim 1, wherein each monomer of said dimeric FAP α molecule consists of the amino acid sequence of SEQ ID NO: 2.
- 3. The dimeric FAP α molecule of claim 1, produced recombinantly.
- 4. The dimeric FAP α molecule of claim 3, produced by a eukaryotic cell.
- 5. Isolated protein consisting of:
 - (i) the FAP α catalytic domain, and
 - (ii) at least one portion of a non $\mbox{FAP}\alpha$ protein.
- 6. Method for cleaving a terminal dipeptide of formula Xaa-Pro from a molecule, comprising contacting said molecule with a second molecule, said second molecule having FAPα enzymatic activity.

7. The method of claim 6, wherein said second molecule is isolated, dimeric $FAP\alpha$.

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- 8. The method of claim 6, wherein said second molecule comprises an FAPα catalytic domain.
- 9. Method for identifying a substance which interacts with a molecule having FAPα activity, comprising combining said molecule with a sample to be tested, and determining any interaction with said molecule as an indication of a molecule which interacts with a molecule having FAPα activity.
- 10. The method of claim 9, wherein said FAP α molecule is dimeric.
- 11. The method of claim 9, wherein said molecule comprises a $FAP\alpha$ catalytic domain.
- 12. The method of claim 9, wherein said substance is an antagonist of FAPa activity.
- 13. The method of claim 9, wherein said substance is an agonist of FAP α activity.
- 14. The method of claim 9, wherein said substance is an inhibitor of FAP α activity.
- 15. The method of claim 9, comprising combining said substance with a cell extract which has FAPα activity.
- 16. The method of claim 15, wherein said cell extract is an extract of a cell which has been transformed or transfected with a nucleic acid molecule which encodes a molecule with FAPα activity.
- 17. The method of claim 16, wherein said cell is a

prokaryote.

- 18. The method of claim 16, wherein said cell is a eukaryote.
- 19. Method for treating a subject with a pathological condition characterized by abnormal FAPα activity, comprising administering to a subject in need thereof an amount of a substance which interacts with molecules having FAPα activity sufficient to normalize the FAPα activity level in said subject.
- 20. The method of claim 19, comprising administering an inhibitor of FAP α activity.
- 21. The method of claim 20, wherein said inhibitor is a collagen derivative.
- 22. The method of claim 20, wherein said inhibitor is (S) Valyl-pyrrolidine-2(R) -boronic acid.
- 23. The method of claim 19, wherein said substance is an agonist of $FAP\alpha$ activity.
- 24. The method of claim 19, wherein said substance is an antagonist of FAP α activity.
- 25. Method for determining if a substance interacts with a molecule having FAPα activity, comprising combining said substance and said molecule with Ala-Pro-AFC, determining interaction of said molecule with Ala-Pro-AFC, and comparing said interaction to interaction of said molecule with Ala-Pro-AFC in the absence of said substance, wherein a difference therebetween indicate that said substance interactions with said molecule.
- 26. Fusion protein comprising a portion of an FAP α molecule sufficient to retain FAP α activity and a non FAP α amino

- acid sequence, wherein said fusion protein is water soluble.
- 27. The fusion protein of claim 26, wherein said non FAPα amino acid sequence is an amino acid sequence found in a CD8 protein.
- 28. The fusion protein of claim 27, wherein said CD8 protein is a murine protein.
- 29. The fusion protein of claim 27, wherein said CD8 protein is a human protein.
- 30. The fusion protein of claim 27 comprising amino acid 1 to 189 of murine CD8 linked to amino acids 27 through 760 of FAPα.

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FIG. 1

FAP	1	MKTWVKIVFGV*ATSAVLALLVMCIVLRPSRVHNSEENTMRALTLKDILN	49
CD26	1	PW-VLL-LLGAA-LVTIITVPVLNKGTDDATADSRKTYT-Y-K	50
FAP	50	GTFSYKTFFPNWISGQEYLHQSADNNIVLYNIETGQSYTILSNRTMKSV*	98
CD26	51	N-YRL-LYSLRDHYKQ*ELVF-A-Y-N-SVF-E-S-FDEFG	99
FAP	99	*NASNYGLSPDRQFVYLESDYSKLWRYSYTATYYIYDLSNGEFVRGNELP	147
CD26	100	HSIND-SIGILYN-V-QHS-DNKRQLITEERI- fap-1	149
FAP	148	RPIQYLCWSPVGSKLAYVYQNNIYLKQRPGDPPFQITFNGRENKIFNGIP	197
CD26	150	NNT-WVTHWN-DV-IE-NL-SYRWT-K-DI-YT fap-2	199
FAP	198	DWVYEEEMLPTKYALWWSPNGKFLAYAEFNDKDIPVIAYSYYGDE**QYP	245
CD26		VFSAYSTQTEV-L-EF-SSL	249
FAP	246	RTINIPYPKAGAKNPVVRIFIIDT***TYPAYVGPQEVPVPAMIASSDYY	292
CD26	250	K-VRVVT-KF-VVN-DSLSSVTNATSIQITASMLIG-H-	299
FAP	293	FSWLTWVTDERVCLQWLKRV <u>ONVS</u> VLSICDFREDWQTWDCPKTQEHIEES	342
CD26	300	LCDVA-QISR-IYMDYD-SSGR-N-LVARQM-	349
FAP	343	RTGWAGGFFVSRPVFSYDAISYYKIFSDKDGYKHIHYIKDTVENAIQITS	392
CD26	350	TV-R-RP-E-H-TL-GN-FI-NEERC-FQIDKKDCTFK	399
FAP	393	GKWEAINIFRVTQDSLFYSSNEFEEYPGRRNIYRISIGSYPPSKKCVTCH	442
CD26	400	-TV-G-EAL-S-Y-Y-IYKGMGL-K-QLSD-T*KVT-LS-E	448
FAP	443	LRKERCQYYTASFSDYAKYYALVCYGPGIPISTLHDGRTDQEIKILEENK	492
CD26		-NPSVKEQ-R-SL-LYSSVN-KGLRVD-S fap-3	498
FAP	493	ELENALKNIQLPKEEIKKLEVDEITLWYKMILPPQFDRSKKYPLLIQVYG	542
CD26	499	A-DKM-Q-V-M-SKKLDFIILN-TKFQHKLDA	548
FAP		GPCSQSVRSVFAVNWISYLASKEGMVIALVDGRGTAFQGDKLLYAVYRKL	592
CD26	549	KADTRLATT-NIIV-SFSGYIMH-IN-R-	598
FAP	593	GVYEVEDQITAVRKFIEMGFIDEKRIAIWGWSYEIRFITGPCIWNWSFQM	642
CD26	599	-TFE-A-Q-SKV-NGGYVTSMVLGSGSVGFK	648
FAP		WYSSGSSLQLGILRVCLHRE*IHGSPNKDDNLEHYKNSTVMARAEYFRNV	691
CD26	649	CGIAVAPVSRWEYYDSVYT-RYM-L-TPEDRSN-KQ-	698
FAP		DYLLIHGTADDNVHFQNSAQIAKALVNAQVDFQAMWYSDQNHGLSGLSTN	741
CD26	699	ET-EDIASSTAH	748
FAP	742	*HLYTHMTHFLKQCFSLSD	
CD26	749	Q-IP	

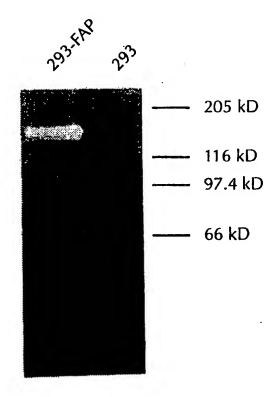
SUBSTITUTE SHEET (RULE 26)

FIG. 2

FAPα	Breast Cancer + A	MFH + C	Healing Wound + E	Renal Cancer G
CD26	<u>—</u> В	D .	÷ F	(+) н

Immunohistochemistry (See Kodachromes)

FIG. 3



SUBSTITUTE SHEET (RULE 26)

INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/04215

IPC(6)	ASSIFICATION OF SUBJECT MATTER ::C07K 14/435; C12N 15/52; A61K 38/43; G01N	33/50; C12Q 1/00					
	:435/4, 7.1, 7.4, 183; 514/2, 12; 530/350 to International Patent Classification (IPC) or to be	th national classification and IPC					
	LDS SEARCHED						
Minimum o	documentation searched (classification system follow	ved by classification symbols)					
1	435/4, 7.1, 7.4, 183; 514/2, 12; 530/350						
Documenta	tion searched other than minimum documentation to	the extent that such documents are included	in the fields searched				
Electronic o	data base consulted during the international search (name of data base and, where practicable	, search terms used)				
APS and	I DIALOG (files 5, 155, 351, 357, and 358) ner, heterodimer						
C. DOC	CUMENTS CONSIDERED TO BE RELEVANT						
Category*	Citation of document, with indication, where	appropriate, of the relevant passages	Relevant to claim No.				
A, P	US 5,587,299 A (RETTIG et al entire document.	24 December 1996, see	1-4 and 19-25				
A	RETTIG, W. J. et al. Regulation and Heteromeric Structure of the Fibroblast Activation Protein in Normal and Transformed Cells of Mesenchymal and Neuroectodermal Origin. Cancer Research. 15 July 1993, Volume 53, pages 3327-3335, especially abstract.						
A	RETTIG, W. J. et al. Cell-surfactions arcomas: Differential expression tissues and cultured cells. Proc. 1988, Volume 85, pages 3110-3	n in normal and malignant Natl. Acad. Sci. USA. May	1-4 and 19-25				
Furthe	er documents are listed in the continuation of Box (C. See patent family annex.					
• Spe	cial categories of cited documents:	"T" Inter document published after the inter	national filing data or priority				
"A" door	mment defining the general state of the art which is not considered a of particular relevance	date and not in conflict with the applicat principle or theory underlying the inve	ion but cited to understand the				
	ier document published on or after the international filing date	"X" document of particular relevance; the considered novel or cannot be consider					
cited	means which may throw doubts on priority claim(s) or which is a to catablish the publication date of another citation or other inl reason (as specified)	when the document is taken alone "Y" document of particular relevance; the					
	ment referring to an oral disclosure, use, exhibition or other	considered to involve an inventive combined with one or more other such being obvious to a person skilled in the	step when the document is documenta, such combination				
"P" docu the p	ament published prior to the interactional filing date but later than priority date claimed	"&" document member of the same patent f	enily				
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Facsimile No		Telephone No. (703) 308-0196	WIT				
Form PCT/IS	A/210 (second sheet)(July 1992)+						

INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/04215

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
Please See Extra Sheet.
As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. X As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.: 1-4 and 19-25
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/04215

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be searched, the appropriate additional search fees must be paid.

Group I, claims 1-4, drawn to FAP-a homodimers.

Group II, claim 5, drawn to a heterodimer.

Group III, claims 6-8, drawn to a method for cleaving.

Group IV, claims 9-18, drawn to a method for identifying substances.

Group V, claims 19-24, drawn to a method of treatment.

Group VI, claim 25, drawn to a method for determining substance interaction.

Group VII, claims 26-29, drawn to a fusion protein.

The inventions listed as Groups I-VII do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons: The special technical feature of Group I is the FAP- α homodimer. None of the other groups requires this homodimer. The other groups require a heterodimer or an unspecified compound with a particular activity. It is noted that PCT Rule 13 does not provide for multiple products and methods within the same application.